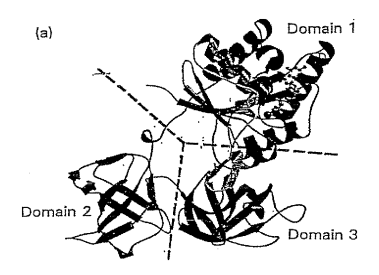
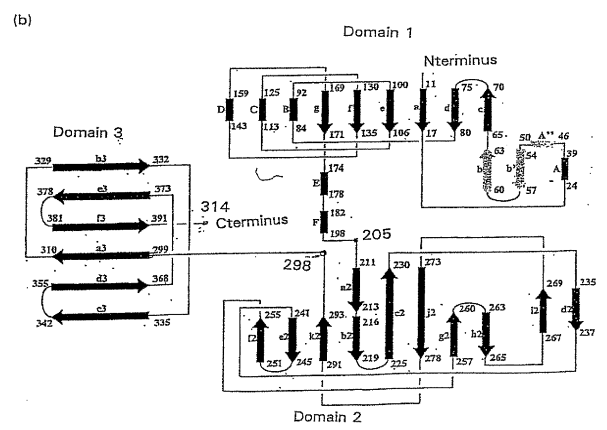
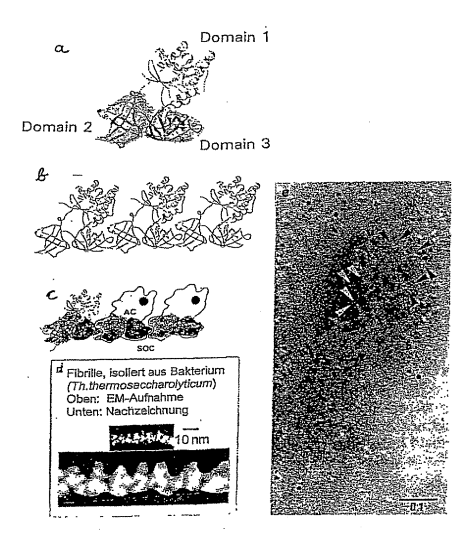
Figur 1





Figur 2



3/15

Fig. 3

(a1)

Vektor pEGFP (Clontech):

CTG CTC GAC ATG TTC ATT TCG CCG GCG CTG AGA TCT TAA GGT

BsrGI-Schnittstelle:

EcoRI-Schnittstelle:

T GTACA G AATTC ACATG T CTTAA G

Synthetisch hergestelltes Oligonukleotid zur Einklonierung des His-Tags in den Vektor:

5'BsrGI BsrGI EcoRI 3'

G TAC AAG CTT CAT CAC CAT CAC CAT CAC TAA CTG TAC AAG TAAG
TTC GAA GTA GTG GTA GTG GTA GTG ATT GAC ATG TTC ATTCTTAA

3'
Tyr-Lys-Leu-His-His-His-His-His-STOP-

Ergebnis: pEGFP(His)

(a2)

Vektor pEGFP(His):

GCC TGC AGG -%- ACC ATG GTG CGG ACG TCC -%- TGG TAC CAC

PstI-Schnittstelle:

NcoI-Schnittstelle:

CTGCA G G ACGTC C CATGG GGTAC C

Fusionsstellen zum EF-Tu-Gen:

Start EF-Tu Start EGFP

5' PstI HindIII Ncol 3'
ACT AGC TGC AGC ATG TCT AAA -%- CTG GGC AAG CTT ACC ATG GTG
TGA TCG ACG TCG TAC AGA TTT -%- GAC CCG TTC GAA TGG TAC CAC

Thr-Ser-Cys-Ser-Met-Ser-Lys-----Leu-Gly-Lys-Leu- Thr-Met-Val

(a3)

Fusionsstellen zur Domäne 3:

5' PstI Cys HindIII Ncol 3'
ACT AGC TGC AGC GCT AAG CCG -%- CTG GGC TGC AAG CTT ACC ATG GTG
TGA TCG ACG TCG CGA TTC GGC -%- GAC CCG ACG TTC GAA TGG TAC CAC
3''
Thr-Ser-Cys-Ser-Ala-Lys-Pro----Leu-Gly-Cys-Lys-Leu-Thr-Met-Val

Sequenz des Konstrukts EF-Tu-GFP-His im Vektor pEGFP (Clontech) (SEQ ID NO:1)

(b1)

pEGFP-Vektor:

NERGINIA CONTROLLA PROGRAMMA A SCAGCIATGA CCATGATTAC GCCAAGCITG AGCGCCCAAI ACGCAAACCG CCICICCCCG CGCGIIGGCC GAIICAIIAA IGCAGCIGGC ACGACAGGII ICCCGACIGG AAAGCGGGCA

EF-Tu:

CATGC GTG G B

AACTGCTGGA CAGGTACTGG CGGCCGTCAT TGCCGGAAGG GACGGTCTGC TCTGTCTCAG AGTGGGAAGC TTCCTGCTGC CAAAGTTGGT GATAACGCGC CGCACACGTA TGGTAGTTGC ATCATCGTGT GTAAAACTAC ACCAT GGCGACGCAG TGACAAGCCG GCGGTATCAT ATGTTCCGCA CGAACGTGGT AAGATGAAGG CGCGATGGAC GTTGACCACG CGACCAGATC GGCGCGATCC CGTTCCGTAC TTCGTGAACT CCCGTCACTA CGTGTAGAAC TGGCGTTGAA GTGAAGAAAT ATTCTGTCCA CGTGACTGGT TCCACCCGAT AAAGTTCTGG AGCGTGCTAT GAAATGGAAG AGCGCTGGAA GACACCCCGA TCAGATGGAC GTCAGGTAGG TATCGGCCAC CICGIGCALL CGTTGTAGCT GGTATCAAAC TGAAGTGTAC GTACTACTGA GTTACCCTGA CIGCICTGAA CCGGAACCAG TGTTACCGGT CTACCTGTAC GGAACTGGTT AAAAATTTGA ACGTACAAAA CCGCACGTTA ACGTTGGTAC CCGGTGCTGC CTGCTGGGTC GGCGGTGCTG CGTTGAATA CAMAATGGTT TTGGCGCGGG GIGGIACCGI AGTICGAAIC TICIACTICC AACATGATCA AAGAGCTGCT GTTCGTGGTT TTCTTARATT ACTCAGAAGT TCTGCTGCGT TGAGCACATC TAAAACCTAC ATCACCATCA ACACTICICA CCGCACACCA CCGTCCGCAG GCGACAACAT GGCCGTACCG TATCAAAGAG CCGTACTGGC CTATGTTAAA CGCAGACTCG GTTGATGACG ATCTCCGGTC ACGIAGGIGI CACTCCGATC GCTTCCTGGA GCTGGTGAGA CACCATCAAG GINAIGCCGG CCGTGAAGGC TCAAAGGCTA CGGGCGACGA GAACTGGCTG CGTATTCTCC AAATCGTTGG GGCCCGATGC ATGCGACATG GCAATCACCA AGCTCGTGGT GGCACGCCGA ACTCCGTTCT CGTAGAGATG GTTTCGCAAT GAAGAAGTTG CGAAGGCCGT CTAAGCCGGG TCCTGAACAA CGATCGAAGA ATGTCTAAAG TACGACTICC SAAAATCCTG CGGAAGAAAA GACTGCCCGG TGCGACTGAC TCTGACCGCT

pEGFP-Vektor: AAGCTTA關

0.1

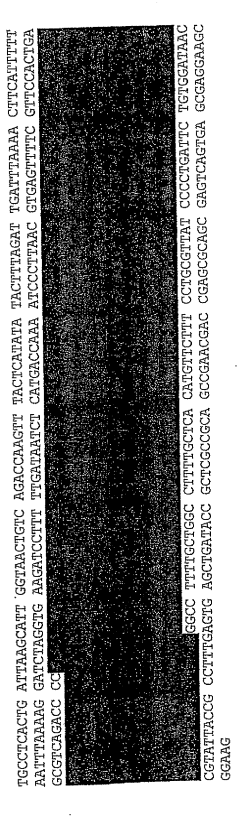
MINION COACTION AGGCCAAGGA GCIGIICACC GGGGIGGIGC CCAICCIGGI CGAGCIGGAC GGCGACGIAA ACGGCCACAA

TTCAAGGAGG AGCAGAACAC CCACATGAAG GCAGAAGAAC ACGGCAACTA ACCGGCAAGC TGGACGAGCINGRIGINA A GACCACTACC CCTGAGCAAA CATCTGCACC GGCCATCGAC TGGCCGACAA GCTACCCCGA TTCAAGGACG CCCAGICCGC ACTCTCGGCA GCAGCTCGCC CACCATCTTC TCGAGCTGAA CCCTGAAGTT TGCTTCAGCC GTCTATATCA CGCCGGGATC TACCTGAGCA CAGCCACAAC ACGGCAGCGT CGGCGTGCAG GTGAACCGCA GGCAAGCTGA PCCAGGAGCG AACATCGAGG TCGTGACCGC CGACAACCAC TGCCACCTAC CCCTGACCTA GAAGGCTACG CGACACCCTG ACAACTACAA GATCCGCCAC CIGCIGGAGT TGCTGCTGCC AGTTCGAGGG AAGCTGGAGT GCGAGGGCGA CTCGTGACCA CGCCATGCCC GACGGCCCCG TCACATGGTC TCTTCAAGTC GCCGAGGTGA CCTGGGGCAC TGAACTTCAA TCCGGCGAGG CTGGCCCACC AGAAGCGCGA STICAGCGIG GGCATCAAGG CCCCATCGGC CAGCACGACT ACGCCAACAT TGCCCGTGCC CAAGACCCGC

# His-Tag: CTTCATCACC ATCACCATCA CTAACTGTAC AAGTAAMENT

CTCATGAGAC CGTCAGGTGG TIGICIGIAA ACTATGCGGC TACCGCATCA TACGGGCCCT AAGGAGAAAA TATGTATCCG TAGICGGCCG CGGTCACAGC GGCTGGCTTA GTTTCTTAGA TACATTCAAA ATAGGCCTAC CGGGTGTCGG ACAGATGCGT GATAATAATG CTCCGGGAGA GAAATACCGC GTCAAAAATA TTAATGTCAT TTTTTTAAA ACACATGCAG CGGGTGTTGG GAGT TATGCGGTGT CTTGTCTGGT AAAACCICIG GGCGCGTCAG TTTTTATAGG TATTGTTTA GARARAGGAR GATGACGGTG GCGGAACCCC CCATTACCAA AGCCCGTCAG GAGTGCACCA GATACGCCTA CAATAATAT GGAAATGTGC ATAAATGCTT ATTGTACTGA GGAGCAGACA AGGGCCTCGT CAACTGAGCG CCGGTCGCTA GCGTTTCGGT DEGFP-Vektor: AATAACCCTG GGCGCCTTA CACTTTTCGG ATCAGAGCAG TICGICICGC GCGGATGCCG

CCTCCCGTA TCGTAGTTAT CTACACGACG GGGAGTCAGG CAACTATGGA TGAACGAAAT AGACAGATCG CTGAGATAGG



Manna Lac-Promotor

m Lac-Operator

Ribosomen-Bindungsstelle

puc Plasmid-Replications-Origin

Ampicillin-Resistenz-Gen

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Die Sequenz enthält vier silent mutations ( on the same of ), die laut Sequenzanalyse eindeutig vorhanden sind:

(1) Soll: TAT, Ist: TA -> Tyr; Codon usage (gesamtes Genom E. coli) ändert sich von 16,2 zu 12,2 (2) Soll: TAC, Ist: TA -> Tyr; Codon usage (gesamtes Genom E. coli) ändert sich von 12,2 zu 16,2 (3) Soll: GCA, Ist: GC -> Ala; Codon usage (gesamtes Genom E. coli) ändert sich von 20,1 zu 33,6 (4) Soll: ATT, Ist: AT -> He; Codon usage (gesamtes Genom E. coli) ändert sich von 30,3 zu 25,1

(Frequenz pro Tausend)

(SEQ ID NO:2) (Clontech) Sequenz des Konstrukts Domäne 3 von EF-Tu-GFP-His im Vektor pEGFP

### pEGFP-Vektor:

CACACIMINA ACAGCIATGA CCATGALTAC GCCAAGCITG AGCGCCCAAT ACGCAAACCG CCICICCCCG CGCGIIGGCC GAIICAITAA IGCAGCIGGC ACGACAGGII ICCCGACIGG AAAGCGGGCA CATGC

## Domäne 3 von EF-Tu:

CTGCCGGAAG CGACGGTCTG GCGGCCGTCA CATTCTGTCC AAAGATGAAG TCGCGATGGA TACCATEGAA CECTEC ACGIGACIGG ATCCACCCGA TAAAGTTCTG CGTACTACTG TGTTACCCTG GCGTTGTAGC CTGAAGTGTA GTTGGCGCGG TCAAAATGGT GCCGCACACC AAGIICGAAI GTTCTACTTC CGTTTCGCAA TCCGTGAAGG CGGCCGTACC ACCGTCCGCA GGCGACAACA GGTAATGCCG GCTAAGCCGG GCACCATCAA TTCAAAGGCT GCGTAGAGAT TACTCCGTTC

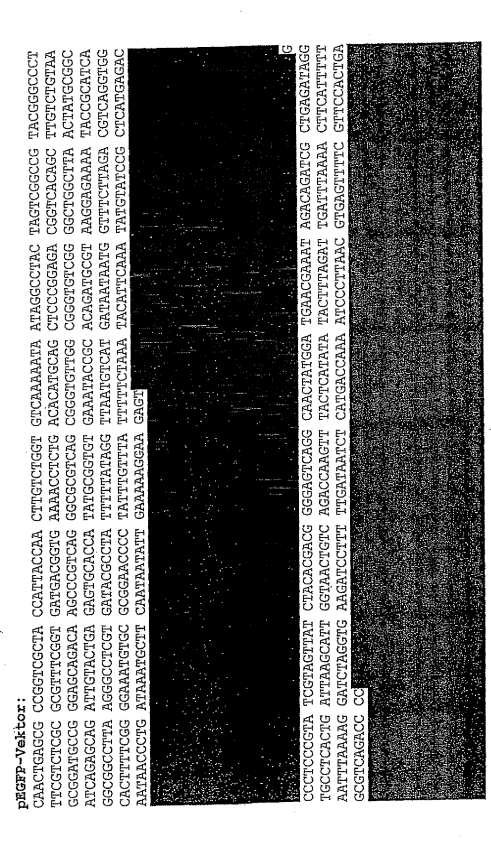
#### pEGFP-Vektor: AAGCTTAB

#### C.

TTCAAGGAGG GCAGAAGAAC AGCAGAACAC CCTGAGCAAA GACCCCAACG CCACATGAAG ACGGCAACTA ACCGGCAAGC GCCGACGTAA ACGGCCACAA TGGACGAGC GACCACTACC CATCTGCACC TTCAAGGACG TGGCCGACAA GCTACCCCGA GGGCATCGAC GCAGCTCGCC CCCAGICCGC ACTCTCGGCA GICIATAICA TCGAGCTGAA CGAGCTGGAC CCCTGAAGTT TGCTTCAGCC CACCATCTTC CGCCGGGATC ACGCCAGCGT CAGCCACAAC TACCTGAGCA CCATCCTGGT CGGCGTGCAG TCCAGGAGCG GTGAACCGCA GGCAAGCTGA AACATCGAGG CGACAACCAC TCGTGACCGC GAAGGCTACG ACAACTACAA CCCTGACCTA CGACACCCTG GGGGTGGTGC TGCCACCTAC CTGCTGGAGT GATCCGCCAC CGCCATGCCC AGTTCGAGGG AAGCTGGAGT TGCTGCTGCC GCTGTTCACC GCGAGGGCGA CTCGTGACCA GACGGCCCCG TCACATGGTC TCTTCAAGTC CCTGGGGCAC TGAACTTCAA AGGCCGAGGA TCCGGCGAGG CIGGCCCACC GCCGAGGTGA GGCATCAAGG AGAAGCGCGA CCCCATCGGC MINGAGCA TGCCCGTGCC CAGCACGACT CAAGACCCGC ACGGCAACAT GTTCAGCGTG

#### His-Tag:

CITCATCACC ATCACCATCA CTAACTGTAC AAGTAA



MANAGE OF THE STATE OF STATES OF STA CGTATTACCG CCTTTGAGTG AGCTGATACC GCTCGCCGCA GCCGAACGAC CGAGCGCAGC GAGTCAGTGA GCGAGGAAGC GGAAG

Bennetal Lac-Promotor

Man Lac-Operator

Ribosomen-Bindungsstelle Ampicillin-Resistenz-Gen

puc Plasmid-Replications-Origin

5. Uniche Pati 6. Grande Neoi 7. Manach Bargi 8. #Aahoe Ecori Die Sequenz enthält eine silent mutation (Meanward), die laut Sequenzanalyse eindeutig vorhanden ist:

Soll: ATT, 1st: ATM -> Ile; Codon usage (gesamtes Genom E. coli) ändert sich von 30,3 zu 25,1 (Frequenz pro Tausend)

